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Journal

Antimicrobial Agents and Chemotherapy, 64(10)

ISSN

0066-4804

Authors

Cuu, Gloria Asua, Victor Tukwasibwe, Stephen <u>et al.</u>

Publication Date

2020-09-21

DOI

10.1128/aac.00721-20

Peer reviewed



Associations between Aminoquinoline Resistance Genotypes and Clinical Presentations of *Plasmodium falciparum* Infection in Uganda

Gloria Cuu,^a ⁽ⁱ⁾Victor Asua,^a Stephen Tukwasibwe,^a Sam L. Nsobya,^{a,b} Ann Nanteza,^c Magambo Phillip Kimuda,^c Arthur Mpimbaza,^{a,d} ⁽ⁱ⁾Philip J. Rosenthal^e

^aInfectious Diseases Research Collaboration, Kampala, Uganda

^bDepartment of Pathology, College of Health Science, Makerere University, Kampala, Uganda

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^cCollege of Veterinary Medicine, Animal Resources and Biosecurity, Makerere University, Kampala, Uganda

^dChild Health and Development Centre, Makerere University College of Health Sciences, Kampala, Uganda

^eDepartment of Medicine, University of California, San Francisco, California, USA

ABSTRACT Mutations that mediate resistance of *Plasmodium falciparum* to aminoquinoline antimalarials are selected by prior drug use and may alter parasite fitness, but associations with clinical presentations are uncertain. We evaluated genotypes in samples from a case-control study of determinants of severe malaria in Ugandan children 4 months to 10 years of age. We studied 274 cases with severe malaria matched by age and geography to 275 uncomplicated malaria controls and 179 asymptomatic parasitemic controls. The overall prevalence of mutations of interest (considering mixed results as mutant) was 67.0% for PfCRT K76T, 8.5% for PfMDR1 N86Y, 71.5% for PfMDR1 Y184F, and 14.7% for PfMDR1 D1246Y. Compared to asymptomatic controls, the odds of mutant PfCRT 76T were lower for uncomplicated (odds ratio, 0.42 [95% confidence interval, 0.24 to 0.72]; P < 0.001) or severe (0.56 [0.32 to 0.97]; P = 0.031) malaria; the odds of mutant PfMDR1 86Y were lower for uncomplicated (0.33 [0.16 to 0.65]; P < 0.001) or severe (0.21 [0.09 to 0.45]; P < 0.001) malaria; and the odds of mutant PfMDR1 1246Y were higher for uncomplicated (1.83 [0.90 to 3.98]; P = 0.076) or severe (2.06 [1.01 to 4.55]; P = 0.033) malaria. The odds of mutant PfMDR1 184F were lower in severe than asymptomatic (0.59 [0.37 to 0.92]; P = 0.016) or uncomplicated (0.61 [0.41 to 0.90]; P = 0.009) malaria. Overall, the PfCRT 76T and PfMDR1 86Y mutations were associated with decreased risk of symptomatic malaria, PfMDR1 1246Y was associated with increased risk of symptomatic malaria, and PfMDR1 184F was associated with decreased risk of severe malaria. These results offer insights into parasite genotypes in children with different presentations, although the basis for the identified associations is likely complex.

KEYWORDS malaria, *Plasmodium falciparum*, drug resistance, aminoquinoline, genotype, drug resistance mechanisms

Malaria, especially falciparum malaria, remains an enormous problem in Africa (1). The treatment of falciparum malaria is challenged by resistance to most available drugs (2). Of particular concern is delayed parasite clearance after treatment with artemisinins and resistance to some artemisinin-based combination therapy (ACT) partner drugs, both of which are now widespread in parts of Southeast Asia, leading to frequent treatment failures (3–6). However, the efficacy of ACTs, including the most widely used regimen, artemether-lumefantrine, appears to remain strong in Africa (7).

Genetic polymorphisms that mediate resistance to several antimalarials and that

Citation Cuu G, Asua V, Tukwasibwe S, Nsobya SL, Nanteza A, Kimuda MP, Mpimbaza A, Rosenthal PJ. 2020. Associations between aminoquinoline resistance genotypes and clinical presentations of *Plasmodium falciparum* infection in Uganda. Antimicrob Agents Chemother 64:e00721-20. https://doi.org/10.1128/AAC.00721-20.

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Address correspondence to Philip J. Rosenthal, philip.rosenthal@ucsf.edu.

Received 14 April 2020 Returned for modification 4 June 2020 Accepted 7 July 2020

Accepted manuscript posted online 13 July 2020

Published 21 September 2020

impact the response to key ACT partner drugs have been common in Africa, but prevalences have been changing (7). Mutations in two Plasmodium falciparum putative drug transporters, PfCRT and PfMDR1, mediate resistance to aminoquinoline antimalarials (8-10). In Africa, the PfCRT 76T mutation, which is the primary mediator of resistance to chloroquine, has been common (7), is selected by therapy with aminoquinolines (11-14), and is selected against by regimens including lumefantrine (12, 14-17). Mutations in PfMDR1, notably the 86Y mutation, are also associated with decreased sensitivity to aminoquinolines (8, 10), selected by therapy with aminoquinolines (12, 14, 18–20), and selected against by regimens including lumefantrine (12, 15-17, 19-21). Coincident with increased use of artemether-lumefantrine and decreased use of chloroquine to treat malaria, the prevalence of the PfCRT 76T, PfMDR1 86Y, and PfMDR1 1246Y mutations has decreased in many countries, including Uganda (7, 22-24). One additional PfMDR1 mutation, 184F, is also common in Africa. This mutation does not have a clear role in mediating drug sensitivity, and it may serve to improve the fitness of parasites with the resistance-mediating PfMDR1 86Y mutation; temporal trends in its prevalence in Uganda have not been straightforward (22-24). Overall, P. falciparum has changed remarkably in Africa in recent years, apparently due to countering selective pressures of different antimalarials and to fitness disadvantages of some mutations.

Resistance-mediating *P. falciparum* mutations may come with some fitness cost. Studies utilizing cultured parasites or animal models have demonstrated fitness advantages for certain PfCRT and PfMDR1 genotypes, although results have been complex (25). Direct demonstration of a fitness advantage for wild-type parasites has come from field studies of the consequences of changes in malaria treatment practices. Discontinuation of chloroquine as the standard antimalarial in Malawi in the 1990s was accompanied by reversion to wild-type sequence at the K76T locus (26) and a return to excellent treatment efficacy for chloroquine (27). Later, across much of Africa, with replacement of chloroquine as the standard antimalarial by artemether-lumefantrine, reversion to wild-type sequences at PfCRT K76T and PfMDR1 N86Y and D1246Y has been seen (7). Loss of mutant parasites after decreased chloroquine use offers evidence of negative fitness consequences of transporter mutations, but in many cases, results were confounded by coincident increased use of lumefantrine, which selects for wild-type parasites.

It is of interest to determine the impacts of drug resistance-mediating P. falciparum polymorphisms on the clinical consequences of malaria. Available studies searching for associations between P. falciparum transporter mutations and severe malaria have been inconsistent, with studies from India (28) and Mali (29), but not Sudan (30) or Gabon (31), showing association between the PfCRT 76T mutation and risk of severe malaria. Considering risk of asymptomatic versus symptomatic infection, in Uganda the prevalence of mutant PfCRT 76T, PfMDR1 86Y, and PfMDR1 1246Y genotypes was higher in children with asymptomatic parasitemia than in those with parasitemia and fever (32). In another Ugandan study considering paired samples from the same subject, the odds of symptomatic malaria were lower for infections with mutant than with wild-type sequence at PfMDR1 N86Y but not PfCRT K76T (33). In these studies, it is difficult to assign causality for parasite polymorphisms in mediating malaria outcomes, as other determinants of outcomes, including parasite density, host genetic polymorphisms, and prior antimalarial drug use, may have varied among study subjects. To better characterize associations between resistance-mediating parasite polymorphisms and malaria outcomes, we evaluated key polymorphisms in samples from a case-control study that included subjects with severe malaria, subjects with uncomplicated malaria, and asymptomatic community controls.

RESULTS

Study populations. Ages of the three study groups were similar, all with median age of \sim 2 years (Table 1). Each group had nearly equal distributions of boys and girls. Mean parasite density was slightly higher in uncomplicated than severe malaria cases,

TABLE 1 Study populations

Variable	Asymptomatic controls $(n = 179)$	Uncomplicated malaria controls ($n = 275$)	Severe malaria cases $(n = 274)$
Vallable	(11 = 179)	CONTROLS (II = 273)	(11 - 274)
Age [yr, median (IQR)]	2.08 (1.22-3.15)	2.12 (1.30-3.14)	1.97 (1.14–2.98)
Gender [n (%) female]	91 (50.8)	132 (48.0)	135 (49.2)
Parasite density [geometric mean (95% CI)]	1,852 (1,080–3,174)	29,240 (24,730–34,560)	25,900 (20,770-32,290)
Gametocytemia, n (%)	21 (11.7)	43 (15.6)	81 (29.5)
Hyperparasitemia (>200,000/µl), n (%)	0	13 (4.7)	31 (11.3)

but hyperparasitemia (>200,000 parasites/ μ l of blood) was more common with severe malaria. Of the 275 asymptomatic controls in the parent study, 179 (65%) had parasitemia detected by blood smear; parasite density in this group was much lower than that in children with symptomatic malaria. Use of at least one medication with antimalarial activity prior to presentation was reported in 44.0% of uncomplicated and 48.9% of severe malaria subjects (Table 2). In both groups the most commonly used antimalarial was artemether-lumefantrine, and 77.2% of those who received antimalarials prior to presentation received an artemisinin-containing regimen. Data on prior drug use were not collected for asymptomatic children.

Prevalence of *P. falciparum* genetic polymorphisms associated with aminoquinoline resistance. We studied the prevalence of four polymorphisms, PfCRT K76T and PfMDR1 N86Y, Y184F, and D1246Y, in all available samples from children with uncomplicated or severe malaria and in samples from asymptomatic children with positive blood smears for *P. falciparum*. Mutations at each of these loci were previously very common in Uganda, but the prevalence of PfCRT 76T and PfMDR1 86Y and 1246Y has decreased, probably due to decreased use of chloroquine and widespread use of artemether-lumefantrine to treat malaria (22–24). Overall, the prevalence of mutations of interest (considering mixed results as mutant) was 67.0% for PfCRT K76T, 8.5% for PfMDR1 N86Y, 71.5% for PfMDR1 Y184F, and 14.7% for PfMDR1 D1246Y.

Comparative odds of genetic polymorphisms with different presentations of malaria. The prevalence of mutations of interest varied among children with different clinical presentations (see Table S1 in the supplemental material). Our primary analysis was unadjusted and considered mixed genotypes as mutant. Compared to asymptomatic controls, the odds of mutant PfCRT 76T were lower for uncomplicated (odds ratio, 0.42 [95% confidence interval, or Cl, 0.24-0.72]; P < 0.001) or severe (0.56 [0.32 to 0.97]; P = 0.031) malaria, the odds of mutant PfMDR1 86Y were lower for uncomplicated (0.33 [0.16 to 0.65]; P < 0.001) or severe (0.21 [0.09 to 0.45]; P < 0.001) malaria, and the odds of mutant PfMDR1 1246Y were higher for uncomplicated (1.83 [0.90 to 3.98], P = 0.076) or severe (2.06 [1.01 to 4.55], P = 0.033) malaria (Table 3). Compared to uncomplicated malaria controls, the odds of mutant PfCRT 76T, PfMDR1 86Y, or PfMDR1 1246Y did not differ from those for severe malaria cases. The odds of mutant PfMDR1 184F were lower

TABLE 2 Antimalarial drug u	use prior to presentation ^a
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	No. (%) for:			
Antimalarial	Uncomplicated malaria controls ($n = 275$)	Severe malaria cases (n = 274)	P value	
Took at least one type of antimalarial	121 (44.0)	134 (48.9)	0.274	
Antimalarial used				
Chloroquine	1 (0.3)	1 (0.3)	1.0	
Artemether-lumefantrine	87 (31.6)	86 (31.3)	0.39	
Oral artemether	2 (0.7)	0	0.21	
Intravenous artesunate	4 (1.4)	23 (8.3)	0.0018	
Oral quinine	5 (1.8)	9 (3.2)	0.59	
Intravenous quinine	2 (0.7)	16 (5.8)	0.0031	
Sulfadoxine-pyrimethamine	3 (1.0)	5 (1.8)	0.73	
Trimethoprim-sulfamethoxazole	27 (9.8)	14 (5.1)	0.019	

^aSome children received more than one antimalarial. P values are based on Fisher's exact test.

	Prevalence [no. positive/total no. (%)]			OR (95% CI); P value			
Genotype	Asymptomatic controls	Uncomplicated malaria controls	Severe malaria cases	Uncomplicated malaria cases vs asymptomatic controls	Severe malaria cases vs uncomplicated malaria controls	Severe malaria cases vs asymptomatic controls	
PfMDR1 N86Y							
Wild type	121/148 (81.7)	255/274 (93.0)	254/266 (95.5)	1	1	1	
Mutant (univariate)	27/148 (18.2)	19/274 (6.9)	12/267 (4.5)	0.33 (0.16–0.65); <0.001	0.63 (0.27-1.40); 0.226	0.21 (0.09–0.45); <0.001	
Mutant (adjusted)				0.65 (0.25–1.70); 0.389	0.62 (0.29–1.36); 0.239	0.23 (0.06–0.760; 0.017	
PfMDR1 Y184F							
Wild type	42/173 (24.3)	68/274 (24.8)	95/271 (35.0)	1	1	1	
Mutant (univariate)	132/173 (75.7)	206/274 (75.2)	176/271 (65.0)	0.97 (0.60–1.54); 0.897	0.61 (0.41-0.90); 0.009	0.59 (0.37–0.92); 0.016	
Mutant (adjusted)				0.95 (0.48–1.86); 0.882	0.61 (0.41–0.91); 0.016	0.68 (0.36–1.28); 0.233	
PfMDR1 D1246Y							
Wild type	122/134 (91.0)	227/268 (84.7)	212/256 (83.1)	1	1	1	
Mutant (univariate)	12/134 (8.9)	41/268 (15.3)	43/255 (16.8)	1.83 (0.90-3.98); 0.076	1.12 (0.68–1.84); 0.626	2.06 (1.01-4.55); 0.033	
Mutant (adjusted)				1.06 (0.45–2.53); 0.880	1.02 (0.62–1.67); 0.936	1.91 (0.86–4.23); 0.108	
PfCRT K76T							
Wild type	24/114 (21.0)	101/261 (38.7)	85/266 (31.9)	1	1	1	
Mutant (univariate)	90/114 (78.9)	160/261 (61.3)	181/266 (68.0)	0.42 (0.24–0.72); <0.001	1.34 (0.92–1.95); 0.105	0.56 (0.32-0.97); 0.031	
Mutant (adjusted)	. ,	. ,		0.50 (0.26–0.95); 0.036	1.31 (0.90–1.92); 0.150	0.62 (0.33-1.18); 0.151	

TABLE 3 Association of	pol	ymorphisms	and	different	malaria	outcomes ^a
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^aOR, odds ratio. Mutants included mixed genotypes. Results are presented based on univariate analyses and after adjustments for each of the other polymorphisms studied, prior use of artemether-lumefantrine (prior use information was not available for asymptomatic controls), and parasite density.

in severe compared to asymptomatic (0.59 [0.37 to 0.92]; P = 0.016) or uncomplicated (0.61 [0.41 to 0.90]; P = 0.009) malaria. The odds of mutant PfMDR1 184F did not differ between uncomplicated malaria and asymptomatic controls.

For each genotype, we considered adjustment for other polymorphisms, prior use of artemether-lumefantrine, and parasite density (Table 3). After these adjustments, the identified associations were similar to unadjusted associations, but odds were generally less pronounced, with loss of some significant associations. These results are consistent with complex associations between genotypes, other factors, and clinical outcomes.

DISCUSSION

We assessed associations between P. falciparum genetic polymorphisms linked to aminoquinoline resistance and clinical presentations in Ugandan children enrolled in a case-control study of severe malaria. The two mutations most clearly associated with aminoquinoline resistance, PfCRT 76T and PfMDR1 86Y, were less common in parasites causing clinical illness than in isolates from children with asymptomatic infections; thus, these mutations were associated with decreased risk of symptomatic malaria. The PfMDR1 1246Y mutation had the opposite association; it was more common in isolates from symptomatic children and, thus, was associated with increased risk of symptomatic malaria. The PfMDR1 184F mutation, which has unclear impacts on drug sensitivity but may play a role in maintaining parasite fitness, was less common in severe compared to uncomplicated or asymptomatic infections and, thus, was associated with decreased risk specifically of severe malaria. Adjustment for other studied polymorphisms, prior use of artemether-lumefantrine, and parasite density decreased some associations, highlighting complex interactions between individual genetic polymorphisms and relevant variables. Our results identify significant associations between transporter mutations and clinical outcomes, but multiple factors account for clinical presentations of African children with malaria, and we cannot conclude that specific genotypes had causal relationships with clinical presentations. Nonetheless, our results offer insights into the likelihood of sensitivity to different antimalarials in parasites associated with different presentations of malaria.

Prior studies have shown varied associations between *P. falciparum* drug resistancemediating polymorphisms and clinical presentations. Considering associations with asymptomatic versus symptomatic disease in Uganda, the prevalence of mutant PfCRT 76T, PfMDR1 86Y, and PfMDR1 1246Y genotypes was higher in children with asymp-

tomatic parasitemia than in those with parasitemia and fever, suggesting that mutant parasites were less likely than the wild type to be associated with symptomatic disease (32). However, in a subsequent study considering paired samples from different episodes in the same subject, to control for potential confounders, differences in prevalences of mutant genotypes between asymptomatic and symptomatic disease were less pronounced and significant only for the PfMDR1 86Y mutation (33). These results highlight the challenge of defining causal associations between specific genotypes and clinical presentations. Potential confounders include the following. First, children presenting with symptomatic disease presumably were more likely to have had recent prior episodes of malaria and, thus, were more likely to have recently received antimalarial therapy with artemether-lumefantrine, which selects for wild-type genotypes. Second, those with symptomatic disease had higher parasite densities and, thus, higher multiplicities of infection than those who were asymptomatic, increasing the likelihood of identifying multiple genotypes, including wild-type isolates, in symptomatic children. Our new study again showed associations between the PfCRT 76T and PfMDR1 86Y mutations and decreased risk of symptomatic infection. Interestingly, the PfMDR1 1246Y mutation had the opposite result; it was associated with increased risk of symptomatic infection, although associations for this polymorphism were modest and not significant after adjustments for other genotypes and prior drug use. This result was unexpected and is not easily explained. As in prior studies, our case-control design did not allow us to control for all potential confounders, but our results show clear differences in parasites seen in symptomatic compared to asymptomatic infections.

Considering genotypes of parasites causing uncomplicated versus severe malaria, potential increased virulence of wild-type parasites might be balanced by risks of progression to severe malaria in those with mutant parasites unresponsive to initial therapy, in particular in older studies in which the standard therapy was chloroquine (28-31). We gathered information on prior drug therapy at the time of study enrollment. Prior use of chloroquine or other aminoquinolines was rare. Prior use of artemether-lumefantrine was common and of similar prevalence in isolates from children presenting with uncomplicated or severe malaria. Unfortunately, we did not collect data on prior drug use from asymptomatic controls, but it is anticipated that recent drug use was uncommon in those who were clinically well. Thus, the selective pressure of recent use of artemether-lumefantrine may explain, at least in part, differences in genotypes between those with symptomatic and asymptomatic parasitemia. Only one studied polymorphism, PfMDR1 Y184F, showed an association with severe malaria: 184F mutant parasites were associated with decreased risk of severe compared to uncomplicated or asymptomatic infections. Interestingly, the PfMDR1 184F mutation, which is common across Africa, may play a role in maintaining the fitness of resistant parasites. In another new study characterizing relative in vitro fitness of parasites with different genotypes, the 184F mutation appeared to stabilize parasites with the 86Y mutation, but in parasites with the wild-type N86 sequence, the wild-type Y184 genotype conferred improved fitness (unpublished data). Although fitness and virulence are distinct properties, these in vitro results are consistent with the conclusion that parasites with the wild-type PfMDR1 Y184 genotype are more likely than mutant parasites to be associated with severe malaria.

Multiple factors influence the clinical outcomes of *P. falciparum* infections, including inherent parasite fitness, host immunity, multiplicity of infection, drug pressure over the course of infection, and host genetics, and it is difficult to determine whether impacts of genotypes on parasite fitness or virulence directly influenced clinical outcomes in our study children. Nonetheless, our results offer insight into the genotypes of parasites in children with different clinical presentations. Notably, children presenting with uncomplicated or severe malaria were more likely to harbor PfCRT K76 and PfMDR1 N86 wild-type parasites that are typically highly sensitive to aminoquinolines but also less sensitive than mutant parasites to artemether-lumefantrine, the first-line therapy for uncomplicated malaria in Uganda and most African countries. Our current understanding of drug resistance in Africa suggests that, even in the setting of PfCRT/PfMDR1

wild-type parasites, artemether-lumefantrine is highly efficacious for the treatment of uncomplicated malaria, but, as has been the case with other artemisinin-based combinations, some loss of activity of lumefantrine might foretell decreasing efficacy of artemether-lumefantrine to treat symptomatic malaria in Africa.

Our study had important limitations. First, any case-control study is necessarily limited by imperfect matching, although in our case efforts were made to match all three groups by age and place of residence, limiting impacts of varied immunity on results. Second, consideration of multiple comparisons limited available sample size and, thus, statistical power for some comparisons. Third, our decision to classify mixed isolates as mutant in all comparisons may have introduced bias for some analyses, but we chose this approach due to its clarity and simplicity. Fourth, information on prior drug use was not obtained for asymptomatic individuals. We assume that asymptomatic children were much less likely than those with symptomatic infections to have used antimalarials shortly before presentation and study enrollment, and decreased selective pressure of AL may well explain the increased prevalence of PfMDR1 86Y and PfCRT 76T mutant genotypes in this group. Fifth, two important polymorphisms, PfMDR1 N86Y and PfMDR1 D1246Y, were quite uncommon, after a remarkable decrease in prevalence in recent years (23, 24), limiting sample size for some comparisons. Finally, we studied only a small set of parasite polymorphisms; full impacts of diverse parasite sequences on clinical outcomes of *P. falciparum* infection were beyond the scope of our study.

In summary, in a case-control study in Uganda, the PfCRT 76T and PfMDR1 86Y mutations were associated with decreased risk of symptomatic *P. falciparum* infections, and the PfMDR1 184F mutation was specifically associated with decreased risk of severe malaria. It is not clear if the studied mutations had causal relationships with clinical presentations, but our results offer insight into genotypes of parasites in children presenting with symptomatic and severe malaria in Africa, including the possibility that parasites from those with symptomatic disease are less responsive to standard therapy with artemether-lumefantrine than are parasites from those with asymptomatic infection.

MATERIALS AND METHODS

Study design. A matched case-control study was conducted in Jinja District, Uganda, in 2015 to 2016 to identify determinants of severe malaria in Ugandan children (34, 35). Cases, recruited from the Children's Ward of Jinja Regional Referral Hospital, were children aged 4 months to 10 years with severe malaria, defined as a positive malaria blood smear (>2,500 parasites/ μ l) and severe anemia (hemoglobin, <5g/dl), impaired consciousness (Blantyre coma score, <4), or respiratory distress (intercostal or subcostal recession without crackles or other evidence of pneumonia upon auscultation). Two types of controls were selected for each case, both recruited from the Busoga Subregion, the catchment area of the Jinja Regional Referral Hospital. First, uncomplicated malaria controls, recruited from level III or IV health centers, were defined as children with fever and a blood smear positive for P. falciparum parasites $(>2.500 \text{ parasites}/\mu)$ but without clinical evidence of severe malaria, danger signs (36), or other known causes of febrile illness. Second, asymptomatic controls were defined as children, recruited from the community, without illness or a history of fever in the 2 weeks prior to enrollment. Both control groups were matched to village and date (within 1 month) of a case. Target age groupings for matching were 6 to <12 months, 1 to <3 years, 3 to <5 years, and \geq 5 years. Caregivers of children with severe or uncomplicated malaria were asked to detail all medicines received during the acute illness, with specific drugs verified using physical descriptions, observations of medicines, and/or prescriptions.

Ethics. Institutional approval was obtained from the Uganda National Council of Science and Technology and the Institutional Review Boards of the College of Health Sciences, Makerere University, and the University of California, San Francisco. Written informed consent was obtained from the parent or guardian of each study subject.

Malaria blood smears. Thick blood smears were stained with 10% Giemsa for 10 min and initially read by health facility staff as part of routine care. The eligibility of severe malaria cases and uncomplicated malaria controls was confirmed upon ascertainment of a positive malaria smear and adequacy of parasite density by a study laboratory technician. Thick blood smears reported as positive were sent to a central laboratory to be read by two expert microscopists for confirmation and determination of parasite densities. If reads were discordant (>25% in parasite density), a third read was performed.

Characterization of *P. falciparum* **genetic polymorphisms.** Parasite DNA was extracted from dried blood spots from samples positive by microscopy using Chelex (37). Sequences of *pfcrt* and *pfmdr1* alleles of interest were determined using a ligase detection reaction-fluorescent microsphere assay, as described previously (38), with minor modifications, including nested PCR amplification of templates, also as described previously (39).

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Analysis. Data were entered using Microsoft Access and analyzed using STATA (version 14). Descriptive statistics were reported as proportions and medians with interquartile ranges. Baseline characteristics in each study group were compared using the chi-square test and the Wilcoxon signed rank test for categorical and continuous data, respectively. Associations between drug resistance-mediating *P. falciparum* polymorphisms and malaria outcomes were initially assessed using cross-tabulations with chi-square tests. To account for confounding and interaction between different polymorphisms, logistic regression was used to measure independent associations between each polymorphism (with adjustment for other polymorphisms, prior use of artemether-lumefantrine, and parasite density) and the clinical outcome groups.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only. **SUPPLEMENTAL FILE 1**, PDF file, 0.02 MB.

ACKNOWLEDGMENTS

This study was supported by funding from the National Institutes of Health (Al075045, Al089674, TW009343, and TW007375).

We thank health workers at the Children's Ward of Jinja Regional Referral Hospital and other participating health facilities for their contributions; Deborah Ekusai Sebatta for administrative support; Simon Peter Kigozi, Thomas Katairo, and Patience Nayebare for methodological support; and study subjects and their parents, guardians, and caretakers for participation in the study.

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